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Specification and Drawings, as originally plication for Patent Serial No: 2,219,713, on October 29, 1997, by McGILL UNIVERSITY, assignee of Philippe Séguéla and Kazimierz Babinski, for Dna Pacoding a Human Proton-Gated Ion Cl ding a Human Proton-Gated Ion Channel and Uses Thereof

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November 16, 1998

Date

# ABSTRACT OF THE INVENTION

The present invention relates to a novel DNA sequence encoding a novel subtype of human proton-gated channel (ASIC3); and uses of the sequence thereof.

# DNA ENCODING A HUMAN PROTON-GATED ION CHANNEL AND USES THEREOF

### BACKGROUND OF THE INVENTION

### (a) Field of the Invention

The invention relates to a DNA sequence encoding a novel subtype of human proton-gated channel; and uses of the sequence thereof.

### (b) <u>Description of Prior Art</u>

The neuronal excitation induced by the contact 10 of acid on peripheral nerve endings has been linked to the activation of specific proton-sensitive cation channels expressed in primary sensory neurons of mammals (Rang et al. (1991) Br. Med. Bull. 47:534-548). The prolonged pain associated with the contact of acid 15 on peripheral nerve endings is due to the activation of non-inactivating proton-gated channels. The duration of the acid-induced pain could neither be explained by the properties of the proton-gated channel ASIC1 cloned from rat (Waldmann et al. (1997) Nature 386:173-177) 20 and human (Garcia-Anoveros et al. (1997) Proc. Ntal. Acad. Sci. (USA) 94:1459-1464) central neurons, nor by the properties of the proton-gated channel ASIC2 cloned also from rat (Waldmann et al. (1997) Nature 386:173-177) and human (Price et al. (1996) J. Biol. Chem. 25 271:7879-7882) central neurons. ASIC1 is sensitive to pH 6.5 and lower but inactivates Waldmann et al. (1997) Nature 386:173-177). ASIC2 is sensitive to pH lower than 6 and inactivates rapidly.

It would be highly desirable to be provided with the primary structure of non-inactivating protonactivated channels from human sensory neurons and means for their functional expression.

### SUMMARY OF THE INVENTION

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One aim of the present invention is to provide the primary structure and functional expression of a subtype of non-inactivating proton-gated channel from human sensory neurons.

Another aim of the present invention is to provide a DNA sequence encoding a novel subtype of human proton-gated channel.

In accordance with the present invention there is provided an isolated nucleic acid molecule which consists essentially of the nucleotide sequence depicted in Figs. 1A and 1B.

The isolated nucleic acid molecule of the present invention encode a peptide consisting essentially of the amino acid sequence depicted in Figs. 1A and 1B.

In accordance with the present invention there is provided a vector, preferably an expression vector, selected from the group consisting of plasmids, phage, retrovirus, baculovirus and integration elements, which include the isolated nucleic acid molecule of the present invention.

In accordance with the present invention there is provided an isolated nucleic acid molecule, which is capable of hybridizing to the isolated nucleic acid molecule depicted in Figs. 1A and 1B, wherein the hybridization occurs at about 35°C to about 65°C and in 5X SSPC and 50% formamide or equivalent hybridization conditions thereto.

In accordance with the present invention there is provided a method of using the isolated nucleic acid molecule depicted in Figs. 1A and 1B, or a sequence which hybridizes under stringent condition to the sequence depicted in Figs. 1A and 1B, to produce a peptide consisting essentially of the amino acid sequence

depicted in Figs. 1A and 1B, which comprises the steps of:

- a) transforming a host with a DNA sequence capable of encoding the peptide;
- b) incubating the host under conditions which allows the sequence to be express;
  - c) isolating the peptide from the host; and
  - d) recording or imaging the activity of the peptide from the host.
- The preferred host is selected from the group consisting of bacteria, yeast, fungi, mammalian cells, plant cells, and insect cells.

In accordance with the present invention there is provided a method of using the peptide encoded by the amino acid sequence depicted in Figs. 1A and 1B or domains of the peptide, to produce antibodies, which comprises the steps of:

- a) immunizing a host with the peptide or domains of the peptide for a time sufficient for an immunogenic reaction to occur; and
  - b) isolating antibodies from the immunized host.

### BRIEF DESCRIPTION OF THE DRAWINGS

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Figs. 1A and 1B illustrate the primary struc-25 ture of the cDNA (1732 bases) encoding the full-length human ASIC3 (hASIC3) channel subunit. The coding region of 531 amino acids encoded in the mRNA corresponds to nucleotides 22 to 1614;

Fig. 2 illustrates the recording of non-inactivating cationic current induced by strong acid (pH 4.0) in *Xenopus* oocytes injected with hASIC3 clone alone in pcDNA3 vector; and

Fig. 3 illustrates the recording of non-inactivating cationic current induced by weak acid (pH 6.5)

in Xenopus oocytes co-injected with hASIC3 clone and rat P2X2 clone both in pCDNA3 vector.

# DETAILED DESCRIPTION OF THE INVENTION

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# Molecular cloning of hASIC3 and in vitro translation

Using the TBLASTN algorithm (Altschul et al. (1990) J. Mol. Biol. 215:403-410), virtual screening of conserved domain database with the dbEST LXTFPAVTLCNXN of ASIC1 and ASIC2 subunits led to the identification of two human fetal brain EST sequences coding for a novel proton-gated channel subunit (EST IDs # AA449579 and AA429417). The clone tagged by EST #AA449579 was sequenced on both strands and was shown to encode a full-length human proton-gated channel subunit (Figs. 1A and 1B). Characteristic natural and unique restriction sites for ClaI, SmaI, SacI, NcoI, XhoI and XbaI are indicated by arrowheads.

This hASIC3 clone was transferred into the of eukaryotic vector pcDNA3 . 20 HindIII-NotI sites (Invitrogen) for CMV-driven heterologous expression in HEK-293 cells and Xenopus oocytes. Supercoiled hASIC3 plasmid was used for in vitro translation using the TnT system (Promega) with T7 RNA polymerase and [35S]-Cysteine according to manufacturer's specifications. The apparent molecular weight of monomeric hASIC3 subunits was 57±3 kiloDaltons, in excellent agreement with the molecular weight of 58.8 kiloDaltons calculated from the predicted primary sequence of the clone.

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### Functional expression of hASIC3 in Xenopus oocytes

Oocytes surgically removed from mature Xenopus laevis frogs were treated 2 hrs at room temperature with type II collagenase (Gibco-BRL) in Barth's solution under agitation. Selected stage IV-V oocytes were defolliculated manually before nuclear microinjection

(Séguéla et al. (1996) J. Neurosci. 16:448-455) of 10 ng cDNA of hASIC3 in pcDNA3 vector. After 2-4 days of expression at 19°C in Barth's solution containing 10μg/ml gentamycin, oocytes were recorded in twoelectrode voltage-clamp configuration using a OC-725B amplifier (Warner Inst.). Signals were acquired and digitized at 500 Hz using a Macintosh IIci equipped with an A/D card NB-MIO16XL (National Instruments) then traces were post-filtered at 100 Hz in Axograph (Axon Acidic solutions titered Instruments). temperature in Ringer's solution containing 115 mM NaCl, 2.5 mM KCl, 1.8 mM CaCl<sub>2</sub> in 10 mM HEPES were applied during 10 seconds on oocytes by perfusion in constant flow (10 ml/min). During recording, oocyte membrane was clamped at Vh=-100 mV.

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There is shown in Fig. 2 the recording of non-inactivating cationic current induced by strong acid (pH 4.0) in Xenopus oocytes injected with hASIC3 clone alone in pcDNA3 vector. These data demonstrate that hASIC3 alone can associate in functional homomeric cation channels.

There is shown in Fig. 3 the recording of non-inactivating cationic current induced by weak acid (pH 6.5) in Xenopus oocytes co-injected with hASIC3 clone and rat P2X2 clone both in pCDNA3 vector. These data demonstrate that the co-expression of hASIC3 and rat P2X2 changes the pH sensitivity of homomeric hASIC3 or leads to the formation of heteromeric pH-sensitive channels.

The present invention will be more readily understood by referring to the following examples which are given to illustrate the invention rather than to limit its scope.

#### EXAMPLE I

# Functional expression of recombinant ASIC3 channels in eukaryotic c lls

Development of analyseic therapeutical compounds used for the clinically-relevant pharmacological modulation, inhibition or activation of human ASIC3 channels and homologous receptors.

## 10 EXAMPLE II

# Uses of antibodies directed against human ASIC3 channel subunits

Polyclonal or monoclonal antibodies can be directed against a bacterial fusion protein containing predicted antigenic domains of hASIC3 subunit, or can be directed against peptides from the predicted amino acid sequence of hASIC3 subunit.

#### Potential uses:

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20 Regional and cellular in situ immunolocalization of mammalian ASIC3 channels in cells naturally or artificially expressing ASIC3 channels.

Immunoprecipitation of mammalian ASIC3 channels for purification of ASIC3 channels and associated proteins, quantitation of ASIC3 channels and associated proteins.

Western blot detection of mammalian ASIC3 channels from cells naturally or artificially expressing ASIC3 channels.

30 Identification of members of the mammalian ASIC gene family using antibodies for screening expression cDNA libraries.

# EXAMPLE III

# Uses of human ASIC3 DNA sequenc

the novel members of Identification of mammalian ASIC channel family as potential therapeutic targets using hASIC3 channel subunit sequence for the design of nucleic acid hybridization probe or PCR the degenerate oligonucleotide primers. While invention has been described in connection with specific embodiments thereof, it will be understood 10 that it is capable of further modifications and this application is intended to cover any variations, uses, or adaptations of the invention following, in general, the principles of the invention and including such departures from the present disclosure as come within 15 known or customary practice within the art to which the invention pertains and as may be applied to the essential features hereinbefore set forth, and as follows in the scope of the appended claims.

### WE CLAIM:

- 1. An isolated nucleic acid molecule encoding peptides consisting essentially of the amino acid sequences depicted in Figs. 1A and 1B.
- 2. The isolated nucleic acid of claim 1, wherein said sequence consists essentially of the nucleotide sequence depicted in Figs. 1A and 1B.
- 3. The isolated nucleic acid of claim 1 or 2, wherein said sequence further comprises a vector selected from the group consisting of plasmids, phages, virus and integration elements.
- 4. The isolated nucleic acid of claim 3, wherein said vector is an expression vector.
- 5. An isolated nucleic acid molecule, which is capable of hybridizing to the isolated nucleic acid molecule of claim 1 or 2, wherein said hybridization occurs at about 35°C to about 65°C and in 5X SSPC and 50% formamide or equivalent hybridization conditions thereto.
- 6. A method of using the isolated nucleic acid molecule depicted in Figs. 1A and 1B, or a sequence which hybridizes under stringent condition to said sequence depicted in Figs. 1A and 1B, to produce peptides consisting essentially of the amino acid sequences depicted in Figs. 1A and 1B, which comprises the steps of:
  - a) transforming a host with a DNA sequence capable of encoding said peptide;

- b) incubating said host under conditions which allows said sequence to be express;
- c) isolating said peptide from said host; and
- d) recording or imaging the activity of said peptide from said host.
- 7. The method of claim 6, wherein said host is selected from the group consisting of bacteria, yeast, fungi, mammalian cells, plant cells, and insect cells.
- 8. A method of using the peptide encoded by the amino acid sequence depicted in Figs. 1A and 1B or domains of said peptide, to produce antibodies, which comprises the steps of:
  - a) immunizing a host with said peptide or domains of said peptide for a time sufficient for an immunogenic reaction to occur; and
  - b) isolating antibodies from said immunized host.

# numan ASIC3

TCCCACGACG	CGGTTCTGGC	CATGAAGCCC A	ACCTCAGGCC	CAGAGGAGGC	CCGGCGGCAG	60
			r s g P	EEA	R R Q	13
	•	*		roGluGluAl	aArgArgGln	
CCCTCCCACA	тессетстт	CGCCAGCAAC 1	TGCTCGATGC	ACGGGCTGGG	CCACGTCTTC	120
P S D I		A S N			H V F	33
		eAlaSerAsn (				
		GCGCCGGGGG 2				180
G P G S	L S L		A A W M		L S V	53
		uArgArgGly 1	MetTrpAlaA	laAlaValVa	lLeuSerVal	
		GGCTGAGAGG (				240
A T F L			V R Y Y		н н Q	73
		lAlaGluArg '	ValArgTyrT	yrArgGluPh	eHisHisGln	
ACTGCCCTGG	ATGAGCGAGA	AAGCCACCGG (	CTCGTCTTCC	CGGCTGTCAC	CCTGTGCAAC	300
TALD	ERE	SHR	L V F P	A V T	L C N	93
ThrAlaLeuA	spGluArgG1	uSerHisArg !	<b>LeuVal</b> PheP	roAlaValTh	rLeuCysAsn	
ATCAACCCAC	TGCGCCGCTC	GCGCCTAACG	CCCAACGACC	TGCACTGGGC	TGGGTCTGCG	360
I N P L			PNDL		G S A	113
IleAsnProL	euArgArgSe	rArgLeuThr	ProAsnAspL	euHisTrpAl	aGlySerAla	
CTGCTGGGCC	TGGATCCCGC	AGAGCACGCC	GCCTTCCTGC	GCGCCCTGGG	CCGGCCCCCT	420
L L G L			AFLR		RPP	133
LeuLeuGlyL	euAspProAl	aGluHisAla :	AlaPheLeuA	rgAlaLeuGl	yArgProPro	
GCACCGCCCG	GCTTCATGCC	CAGTCCCACC	TTTGACATGG	CGCAACTCTA	TGCCCGTGCT	480
A P P G	_		F D M A		ARA	153
AlaProProG	lyPheMetPr	oSerProThr	PheAspMetA	laGlnLeuTy	rAlaArgAla	
GGGCACTCCC	TGGATGACAT	GCTGCTGGAC	TGTCGCTTCC	GTGGCCAACC	TTGTGGGCCT	540
G H S L			CRFR			173
GlyHisSerL	euAspAspMe	tLeuLeuAsp	CysArgPheA	rgGlyGlnPr	oCysGlyPro	
GAGAACTTCA	CCACGATCTT	CACCCGGATG	GGAAAGTGCT	ACACATTTAA	CTCTGGCGCT	600
E N F T	TIF	TRM	G K C Y	T F N	S G A	193
GluAsnPheT	hrThrIlePh	eThrArgMet	GlyLysCysT	yrThrPheAs	nSerGlyAla	
GATGGGGCAG	AGCTGCTCAC	CACTACTAGG	GGTGGCATGG	GCAATGGGCT	GGACATCATG	660
D G A E	LLT	TTR	G G M G	N G L	DIM	213
AspGlyAlaG	luLeuLeuTh	rThrThrArg	GlyGlyMetG	lyAsnGlyLe	uAspIleMet	
CTGGACGTGC	AGCAGGAGGA	ATATCTACCT	GTGTGGAGGG	ACAATGAGGA	GACCCCGTTT	720
L D V Q			V W R D		T P F	233
LeuAspValG	lnGlnGluGl	uTyrLeuPro	ValTrpArgA	spAsnGluGl	uThrProPhe	
					ClaI	
GAGGTGGGGA	TCCGAGTGCA	GATCCACAGC	CAGGAGGAGC	CGCCCATCAT	CGATCAGCTG	780
E V G I			QEEP			253
	_	nIleHisSer			-	
-	Sma				•	
	~			0001001001	00100ma160	0.40
		CTACCAGACC				840 273
		Y Q T				2/3

Fig. 1A

# huma SIC3

		1			
TTCCTGCCAC CGCCCTGGG					900
		SASL		Y E P	293
PheLeuProP roProTrpG	l yAspCysSer	SerAlaSerL	euAsnProAs	nTyrGluPro	
GAGCCCTCTG ATCCCCTAG	G CTCCCCCAGC	CCCAGCCCCA	GCCCTCCCTA	TACCCTTATG	960
		P S P S		T L M	313
GluProSerA spProLeuG	l ySerProSer	ProSerProS	erProProTy	rThrLeuMet	
GGGTGTCGCC TGGCCTGCG					1020
G C R L A C E		VARK		R M V	333
GlyCysArgL euAlaCysG		ValAlaArgL	vsCysGlyCy	sArgMetVal	
TACATGCCAG GCGACGTGC					1080
_	V C S			A H P	353
Y M P G D V P TyrMetProG lyAspValP					
					1140
GCCATAGATG CCATCCTTC					-
AIDAILR				A S T	373
AlaIleAspA laIleLeuA	r gLysAspSer	CysAlaCysP	roasnprocy	salaserini	
	NcoI				
Sa					
	_				
CGCTACGCCA AGGAGCTCT	CATGGTGCGG	ATCCCGAGCC	GCGCCGCCGC		1200
RYAKELS		IPSR			393
ArgTyrAlaL ysGluLeuS					
GCCCGGAAGC TCAACCGCA	G CGAGGCCTAC	ATCGCGGAGA	ACGTGCTGGC	CCTGGACATC	1260
	E A Y	IAEN		LDI	413
AlaArgLysL euAsnArgS	e rGluAlaTyr	IleAlaGluA	snValLeuAl	aLeuAspIle	
TTCTTTGAGG CCCTCAACT	A TGAGACCGTG	GAGCAGAAGA	AGGCCTATGA	GATGTCAGAG	1320
F F E A L N Y				M S E	433
PhePheGluA laLeuAsnT				uMetSerGlu	
CTGCTTGGTG ACATTGGGG					1380
	Q M G				453
LeuLeuGlyA spIleGlyG					
- · · · · · · · · · · · · · · · · · · ·	. 101.11.00011	20101100			
XhoI					
CTCGAGATCC TAGACTACC	r ctgtgaggtg	TTCCGAGACA	AGGTCCTGGG	ATATTTCTGG	1440
LEIL DYL		-		Y F W	473
LeuGluIleL euAspTyrL	e uCysGluVal	PheArgAspL	ysValLeuGl	yTyrPheTrp	
AACCGACAGC ACTCCCAAA	G GCACTCCAGC	ACCAATCTGC	TTCAGGAAGG	GCTGGGCAGC	1500
NRQHSQR				L G S	493
AsnArgGlnH isSerGlnA	r gHisSerSer	ThrAsnLeuL	euGlnGluGl	yLeuGlySer	
CATCGAACCC AAGTTCCCC					1560
HRTQVPH					513
HisArgThrG lnValProH					
		<u>.</u>		XbaI	
				_	
GTCACCAAGA CTCTCTCCG				GCTCTAGACC	1620
V T K T L S A		T C Y L		L.	531
ValThrLysT hrLeuSerA	l aSerHisArg	ThrCysTyrL	euValThrG1	nLeu	
TGCTGTCTGT GTCCTCGGA	CCCCCCCCTG	ACATCCTGGA	CATGCCTAGC	CTGCACGTAG	1680
CTTTTCCGTC TTCACCCCA					1732

Fig. 1B

# Non-desensitizing pH-sensitive inward current in Xenopus Oocytes microinject d with hASIC3

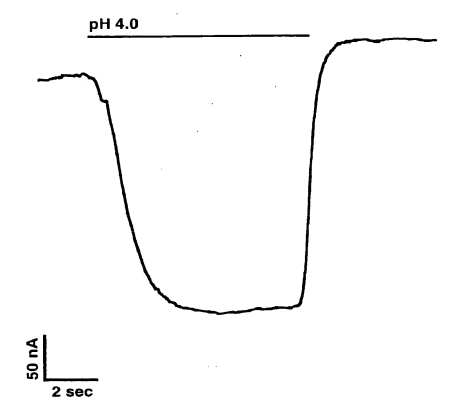


Fig. 2

Non-desensitizing pH-sensitive current in Xenopus oocytes microinjected with human ASIC3 + rat P2X2

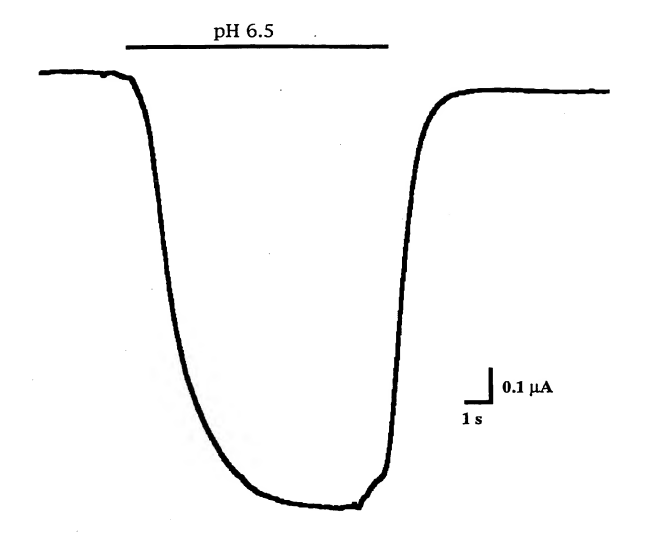


Fig. 3

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